DECLARATION

translation to the best of my knowledge and belief. languages and certify that the following is a true am well acquainted with the English and Japanese that I, the translator of the documents attached, Mameido, Koto-ku, Tokyo, Japan, do solemnly declare I, Hiroji KUMAZAWA, of 9-33-1-308 Dated this /4 May of July, 1992.

PATENT OFFICE

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JAPANESE GOVERNMENT

This is to certify that the annexed is a true copy of the following application as filed with this office.

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Application Number: 18th April, 1991. Patent Application No. 114074/91

Applicant:

TONEN CORPORATION

29th May, 1992.

Matery FUKAZAWA

Seal

Commissioner, Patent Office

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SPECIFICATION

[Title of the Invention]

Process for the Expression of Human Protein Disulfide Isomerase Gene and Process for the Preparation of Polypeptides by Co-Expression with the Gene

[Claims]

- 1. A fusion gene for use in an expression of human protein disulfide isomerase, which is composed of a DNA fragment coding for a human serum albumin prepro sequence and a gene coding for said isomerase.
- 2. A fusion gene for use in an expression of human protein disulfide isomerase, the fusion gene being composed of a DNA fragment coding for a human serum albumin prepro sequence and a gene coding for said isomerase, characterized in that the fusion gene has a base sequence coding for the -24 to +491 amino acid sequence shown in SEQ ID NO:2.
- 3. A fusion gene according to claim 2, wherein said base sequence is the sequence between nucleotide 1 and nucleotide 1545 shown in SEQ ID NO:2.
- 4. A replicable expression vector capable of expressing th fusion gene according to any one of claims 1-3 in a host deli-

- 5. A transformant obtained by transforming a host ceil with the expression vector according to claim 4.
- A transformant according to claim 5, wherein the host cell is a yeast cell.
- 7. A process for the preparation of a recombinant human protein disulfide isomerase, characterized by expressing the fusion gene according to any one of claims 1-3, in the transformant according to claims 5 or 6.
- 8. A process according to claim 7, which comprises the following steps of:

constracting a replicable expression vector capable of expressing the fusion gene according to any one of claims 1-3 in a host cell;

transforming the host cell with said expression vector to obtain a transformant;

culturing said transformant under the conditions capable of expressing said fusion gene, to excrete a recombinant human protein disulfide isomerase; and

recovering said recombinant human protein disulfide isomerase.

9. A process according to claim 8, wherein the recombinant human disulfide isomerase excreted is separated and recovered by means of hydrophobic column chromstography.

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A recombinant human protein disulfide isomarase having an any 491 shown in SEQ 3, the isomerase being obtained by the process according to position from position 1 to amino acid sequence of claims 7-9.

A transformant comprising, in a co-expressible state, the of the claims 1-3 and a foreign be produced. for a polypeptide to to any one fusion gene according gene coding , 64

N A transformant according to claim 11, which is transformed yeast cell 12.

A transformant according to claim 11, wherein said foreign numan serum albumin. #OF gene coding gene is a er,

A process for the preparation of a polypeptide, which comprises the following steps of:

one of claims 11-13 so as to produce comergesing a human protein disuifide isomerase gene and the polypeptide to be produced, in the to any foreign gene coding for transformant according the polypeptide; and

recovering said polypeptide.

A process according to claim 14, wherein said polypaptide is human serum albumin χή (1)

(Detailed Description of the Invention)

(Document Name)	Application For Patent
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(Filing Date)	18th April, 1991.
(Addressed To)	Mr. Satoshi UEMATSU, Commissioner of Patent Office
(Title of Invention)	"PROCESS FOR THE EXPRESSION OF HUMAN PROTEIN DISULFIDE ISOMERASE GENE AND PROCESS FOR THE PREPARATION OF POLYPERTIDES BY CO-EXPRESSION WITH

(Residence/Domicile) (Number of Claims) (Inventor)

THE GENE"

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[Field of the Invention]

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This invention relates to an expression of a gene coding for protein disulfide isomerase which is an enzyme for enhancing the formation of a high-order structure of polypeptide by catalyzing an exchange reaction of a disulfide bond(s) in the polypeptide. The present invention also relates to a co-expression of said gene with a foreign gene coding for a useful polypeptide.

[Prior Art]

revealed the presence of both isomerization reactions of a disulfide bond and of a proline peptide as factors for determining a folding rate of polypeptides (Freedman, Cell, vol.57, pp.1069 - 1072, 1989; Fisher and Schmid, Biochemistry, vol.29, pp.2205 - 2212, 1990). As enzymes which catalyze these slow reactions during the polypeptide folding, peptidyl prolyl cis-trans isomerase (PPI) has been found in the latter case, and protein disulfide isomerase (PPI) and thuse enzymes rise a refolding rate of denatured proteins, thus indicating a possibility of applying them to the in vitro experiments, (Schein, Bio/Technology, vol.7, pp.1141 - 1148, 1989; J. Odaka, Nippon Nogel Kayaku Kaishi, vol.54, pp.1035 - 1038, 1990).

Since PDI is soluble in water and can be isolated relatively easily from the liver of mammals, its properties as a catalyst have been studied in detail. PDI catalyzes the exchange reaction between thiol/disulfide bonds and is capable of undergoing formation,



isomerization or reduction of the disulfide bond in protein substrates (Freedman, Cell, vol.57, pp.1069 - 1072, 1989). It is known that, in vitro, FDI enhances the formation or exchange reaction of a disulfide linkage(s) in molecules of a single domain protein such as RMase and of a multiple domain protein such as serum albumin, or enhances the formation of an intermolecular disulfide bond(s) in a protein having a subunit structure such as immunoglobulin, procollagen or the like (Freedman, Nature, vol.329, p.195, 1987).

ppI from Mammais exists usually as a homodimer of the polypeptide having a molecular weight of about 57,000 and shows a highly acidic pI value (4.2 to 4.3).

The PDI gene from rat liver has been isolated. The amino acid sequence deduced from the DNA sequence of the PDI gene indicated that PDI has an intramolecular duplicate structure consisting of two homologous units. One of these two homologous units has a homology to the amino acid sequence of thioredoxin, indicating that its active site has an amino acid sequence similar to that of thioredoxin (Edman et al., Nature, vol.317, pp.267 - 270, 1985). Thioredoxin enhances the reduction of a disulfide bond in insulin and the exchange reaction of a disulfide bond in RNase in vivo, which indicate that thioredoxin plays a similar role to PDI in the in vivo folding process of proteins (Pigict and Schuster, Froc. Natl. Acad. Sci., USA, vol.83, pp.7643 - 7547, 1985).

Although the amount of PDI present in a living body differs depending on the type of tissues and the differentiation stage of cells, such a difference is correlated with the existence of certain

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In addition, \$01 is localized abundantly in the ranslation of Y-gliadin hardly occurs when an endoplasmic reticulum fraction from which PDI was washed out in advance is used, while the disulfide bond formation is restored by the addition of FDI (Buileid during its secretion. On the basis of these facts, it is assumed olosynthesis of Y-gliadin in a cell-free protein synthesis system through which a protein is known to pass disulfide bond in conjunction with the S that PDI concerns with the formacion of a disulfide bond(s) Such assumption is supported by the results of a study on the ecretory proteins newly synthesized within cells. Freedman, Nature, vol.335, pp.649 - 651, 1988) that the formation of a endoplasmic reticulum secretory proteins.

binding protein that recognizes a signal sequence Asn-X-Ser/Thi addition to the disulfide band formation, PDI concerns with of a peptide to which a sugar chain is bound during N-glycosylation catalycic unit, \$-subunit, of proly1-4-hydroxylase which catalyzes molecular species having partly homologous amino acid sequences to the polyfunctional property of PDI in connection with the protein ydroxylation of proline residues in collagen, to a glycosylation other post-translational modifications of proteins. For example, thyronine binding protein; Cheng et al., J. Biol. Chem., vol. 262, - 11227, 1987), etc. In addition to these facts, some pp.643 - 549, 1987; Geetha-Habib et al., Cell, vol.54, pp.1053 odifications has been suggested on the basis of its homology 1050, 1988), to a thyroid hormone binding protein (trilodo-1process of synthetic protein (Pihlajaniemi et al., 5MBO J., PDI have been found though different from the PDI. 22.11221



certain genadotropic hermones such as fellitropin and lutropin contain amino acid sequences homologous to an amino acid sequence which is regarded as an active site of PDI, and these hormones catalyze the isomerization of a disulfide bend (Beniface et al., Science, vol.247, pp.61 ~ 64, 1990). Also, phospholipase C, an enzyme which hydrolyzes phosphatidylinositol-4,5-bisphosphate into 1,2-diacyl glycerol and inositol-1,4,5-triphosphate, has a domain homologous to PDI in its molecule (Bennett et al., Nature, vol.334, pp.268 - 270, 1988). In consequence, PDI and PDI-like molecules seem to concern in a markedly wide range of vital phenomena, both intracellularly and extracellularly.

has cond(s). In many cases, however, an almost stoichiometric amount of PDI is required to attain an optimum reaction rate. It is expected secretory proteins) in Escherichia coll. Aithough E. coll contains disulfide reductase, the isomerase activity of thioredoxin is low. Although the FDI has extensive functions as described above, cate of a disulfide bond will be slow when a disulfide isomerase thioredoxin which is superior to PDI in terms of the activity as therefore that an intramolecular or intermolecular isomerization main effect of PDI is to form a protein (or a protein aggregate) low activity, and such a slow reaction rate will entail a low aggregates of various eukaryote-originated proteins (especially isomerization of an intramolecular or intermolecular disulfide formation efficiency of a protein having a suitable disulfide bond(s). It is thought that such a low disulfide isomerase is one of the cause of the formation of insoluble molecular having a natural higher-order structure by catalyzing the



On the contrary, since an intramolecular disulfide bond(s) can often be found in secretory proteins, it is thought that a disulfide bond activity resulted from disulfide isomerization is high in cells or tissues which have a high secretion ability. This was indicated strongly by a comparative study on the relative PDI mRNA contents in various rat tissues, in which the contents in organs were found to be liver > pancreas, kidney > lung > spermary, spleen > heart > brain in order (Edman et al., Nature, vol.314, pp.267 - 270, 1985).

environment (compartment) isomerase) should be present which has a high affinity for the large amount of a disulfide-forming enzyme (i.e., disulfide environment (compartment) should be formed in a host cell and a to form a disulfide bond under certain conditions which are suitable proteins) is produced by recombinant DNA techniques, it is necessary its formation. When useful proteins (most of them are secretory different from each other in terms of factors concerning the for each protein to be produced. For this purpose, a suitable formation of a disulfide bond and of an environment which enables prokaryotic cells which have no compartments. Taking this into consideration, prokaryotic cells and eukaryotic cells may be necessary for the suitable folding of a polypeptide will be synthesis system, the formation of a disulfide bond which is inhibited. Such an environment is generated for example in When an environment in a reduced state is given to a protein

The above two points must be considered greatly when a protein having a disulfide bond(s) is produced effectively using recombinant DNA techniques.



However, nothing is in practice known about an in vivo system in which PDI is contained in a large quantity in a suitable compartment where a useful target protein co-exsists, the PDI being capable of acting on the protein.

(Broblems to be solved by the Invention)

In spite of the applicability of PDI to the in vitro refolding of denatured proteins and to the improved productivity of secretary proteins in cells, this enzyme has been prepared only by direct purification from the internal organs. In addition, there are no reports on the interspecific expression of PDI, and on the establishment of a process for its production by means of genetic engineering or a process in which the productivity of a useful polypeptide is improved by the combination of the PDI gene with a gene coding for the polypeptide.

It is objects of the present invention is to provide a fusion gene for use in the expression of PDI, which is composed of a DNA fragment coding for a human serum albumin prepro-sequence and a gene coding for the PDI; an expression vector capable of expessing said fusion gene in a host cell; a cransformant obtained by transforming a host cell with said vector; a process for the preparation of a recombinant human PDI by expessing the fusion gene in said transformant; and the recombinant PDI.

Other objects of the present invention are to provide a transformant containing, in a comexpressible state, both the above mentioned fusion gene and a foreign gene coding for a polypeptide to be produced; and a process for the production of the polypeptide by

co-expressing the human PDI gene and the foreign gene in said cransformant so as to produce the polypeptide.

(Means to solve the Problems)

The present invention has been completed by finding an expression of PDI expression vector integrated a fusion gene for expression of PDI which is composed of a DWA fragment coding for a human serum albumin prepro-sequence and a gene coding for human PDI.

The following describes the present invention in detail:

Clones containing a human PDI cDNA are isolated from the human liver and placenta Agtli cDNA libraries (Clontech, US) by the following procedures:

and placenta Agtli cDNA libraries, after which DNAs from the human liver grown are fixed on a filter. Separately from this, positive clones are screened by hybridization using a 40 mer synthetic oligomer DNA as a probe which corresponds to the complementary strand of a nucleic acid sequence (143-282) of human proline 4-hydroxylase (the p.643, 1987). The phage DNA obtained is digested with EcoRI, and the resultant 150 bp insert DNA is used as a probe for screening the PDI cDNA. Using the probe, the phage DNAs fixed on the filter are screened to isolate positive clones.

Thereafter, a plurality of positive clones obtained in such an anner are digested with EccRI so as to isolate EccRI insert DNA fragments, and a restriction map of the insert of each clone is made. From the comparison of these maps with that reported by

Pihlajaniemi et al., it was estimated that the full length human PDI cDNA was covered by a clone (pappile) from liver and a clone (pappile) from liver and a clone

these clones encoded human FDI cDNA consisting of 2454 base pairs in full length as shown in the SEQ ID No:1. An amino acid sequence deduced from the DNA sequence is also shown in the SEQ ID No:1. In the amino acid sequence, a mature protein seems to be composed of 491 amino acids from Asp¹ to Leu⁴⁹¹, and the 17 amino acid polypeptide preceding Asp¹ seems to be a signal peptide.

According to the present invention, there is provided a fusion gene for use in the expression and production of PDI, which is composed of a DNA fragment coding for a human serum albumin preprosequence and the aforementioned human PDI gene.

As shown in Fig. 1C, the fusion gene is constructed in general by arranging the preprosequence-encoding DNA fragment at the upstream side of the PDI gene. In this instance, however, a leader sequence for transporting human PDI into an appropriate compariment (considered to be endoplasmic reticulum) is not always limited to the BSA prepro-sequence, and other signal sequences or prepro-sequences may also be used as the leader sequence.

More particularly, said fusion gene may be prepared as follows:

The aforementioned clones purplis and purpley DNAs are double—
digested with Scori/PstI and PstI/Bamii, respectively, to produce
DNA fragments of about 490 bp and about 1.3 kbp respectively, the
fragments recovered are ligated with a plasmid vector pUCl19 which
was digested with EcoRI and Bamii to produce phPDIEB in which a Nael

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signal sequence and the PDI sequence on the CDNA by the Nunkel's cleavage site is then introduced into the boundary between the PDI 1.7 kb which does not contain the PDI signal sequence digested with Nael and Hindlil to give a PDI DNN fragment of about method (Kunkel, T.A., Proc. Natl. Acad. Sci., USA, vol.82, p.488 so as to prepare phpDINae, and thereafter the phpDINae is

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resultant digest is ligated with the following whol linker: Separately from this, pUCli9 is digested with EcoRI, and eun

GAGCTCTTAA-5'

S'-AATTCTCGAG

digest is ligated with a prepro-sequence of HSA to produce pUC119Sig which is subsequently digested with Stul and HindIII to give a DNA sequence will be described later in Examples). fragment of about 3,2 kb (A method for synthesizing the HSA prepro-After double-digesting the product with XhoI and BamHI, the

produce phapity; which is in turn digested with EcoRI, blunt-ended kb DNA fragment from pUC119Sig origin are ligated together to sequence (Fig. 2). buman PDI gene is fused to the downstream side of the HSA preprofusion gene in which a leader sequence is modified and in which the with Klanow fragment, and digested with BankI, thereby giving a Thereafter, the 1.7 kb DNA fragment from phPDINae and e tine

are not limited to the above-described techniques, provided that to the present invention and an arrangement of its constituted genes analogs of the inventive fusion gene are not included within the said fusion gene has an ability for expressing PDI. both a process for the preparation of the fusion gene according Although



than human prepared easily from a corresponding gene of any animal origin other scope of the present invention, it is obvious that they can

fusion gene has a DNA sequence coding for the -24 to +491 amino having nuclectide sequences based on the degeneracy of codon, are these instances, all genes which substantially have the same therein, a DNA sequence coding for the +1 to +491 amino acid another embodiment of the present invention, an example of such a function as that of said DNA sequences, for example, derivatives sequence (Aspi-Leu491) shown in SEQ ID NO:1 may also be applied. sequence shown in the SEQ ID NO: 2. fusion gene includes the whole of the base sequence shown in SEQ ID included within the scope of the present invention. According to According to one embodiment of the present invention, said Instead of the PDI gene H

invention in a host cell. vector capable of expressing the fusion gene of the present The present invention also provides a replicable expression

derived from a species compatible with a host to be used, as well as useful vector contains replicon and regulatory sequences which are and have the ability for expressing therein. In general, such a replication origin and a marker sequence which enables selection the present invention thereinto should replicate in a host cell phenotype from transformed cells An expression vector used for the insertion of the linked gene

vector, the plasmid pJDB-ADK-HSA-A (Fig. IC) which has been As a vector for use in the construction of the expression

45 this process, other types of vectors can be used provided that they \sqrt{} esistance gene (Amp^{κ}) and Leu2 gene. The HSA CDWA is removed from resultant fragment is ligated with the aforementioned fusion gene present invention to give the expression plasmid pAHAPDILY1. fragment of about 8 kb thus obtained is dephosphorylated, and the isolosed in Japanese Patent Application Laying-Open (KOKAI) No. The 5'-end of the DWA be used conveniently iehydrogenase I (ADH I) promoter, ADH I terminator, ampicillin blunting with Klenow his plasmid contains HSA cDWA, as well as yeast alcohol mey. capable of expressing the fusion gene fragment, and then digesting with BamHI. chis plasmid by digesting it with Xhoi, 17384 filled by the present applicant

The present invention further provides a transformant obtained by transforming a host cell with the expression vector of the

Examples of such a host include prokaryotes such as E. cali, sacilius subtilis, etc, and yeast. Preferred is a host cell capable of secreting the mature PDI via processing. Preferably, the host cell is yeast such as Saccharomyces cerevisiae, and, in particular, a yeast strain AH22 is suitably used in the case of a preparation of the transformant according to the present invention. It is obvious that eukaryotes other than yeast, for example animal cells, can be used as the host cell, although they are not included within the scope of the invention. Incorporation of the expression vector into a host cell can be carried out easily by conventional means such as calcium chloride, protoplast (or spheroplast) polyethylene glycol, electroporation, etc. When the plasmid phihabbliyl is used

as an expression vector, a desired transformant may be obtained by grown on the plate.

Accordingly, the present invention also provides a process for the preparation of a recombinant human PDI by expressing the fusion gene of the invention in a transformant prepared in the same manner as described above. In one enbodiment of the present invention, the process of the invention comprises the following steps of:

constructing an expression vector which can replicate in a host cell and express the fusion gene of the present invention therein; isolating a host cell transformed with said expression vector; culturing the obtained transforment under such conditions that the fusion gene can be expressed, thereby secreting said recombinant human PDI; and

recovering the recombinant PDI.

If the host is yeast, a human PDI precursor protein is processed to excrete the human recombinant PDI as a gene product. If microorganisms other than yeast, for example E. coli and Bacillus subtilis, are used as host, a non-processed human PDI precursor protein will be obtained.

The recombinant human PDI can be purified easily by separating the transformed cells from a cultured medium by centrifugation, disrupting the cells if necessary, concentrating the supernatant by ultrafiltration or the like, and then subjecting the concentrate to hydrophobic column chromatography. Though not particularly limited, TSK-gel Phenyl-5PW hydrophobic column (Tosoh, Japan) may be used in the chromatography. In this case, the recombinant human PDI may be



eluted by linear gradient of from 0.85 to 0 M ammonium sulfate in borate buffer (pH 8.0) containing NC1 (Fig. 4). It was confirmed that the purified recombinant human PDI has a molecular weight of about 55 kDa based on the SDS-polyacrylamide gel electrophoresis analysis (Fig. 5), and practically has PDI activity as the results of determination of a degree of the refolding of scrambled ribonuclease A (see Examples).

In comparison with the natural type human PDI, it has been found that the recombinant human PDI thus prepared has the same amino acid sequence except that its N-terminal amino acid is changed from Asp to Giy. Accordingly, the present invention provides a recombinant human PDI having the amino acid sequence Gly¹--leu⁴⁹¹ consisting of 491 amino acids shown in SEQ ID NO:3.

The present invention further provides a transformant gene and a computating state.

The fusion gene and the foreign gene in the transformant may be located on the same or different chromosome(s), provided that they are mutually present in a co-expressible state. Transformation of a host cell can be carried out for example by inserting the fusion gene and the foreign gene into the same or different vector(s) and introducing the resulting vector or vectors into the host cell by conventional means such as calcium chloride, protoplast(or spheroplast)-polyethylene glycol, electroporation, etc.

that the polypeptide contains a disultide linkage(s) because the

catalytic effect of an amplified and expressed PDI, that is, acceleration of the formation or exchange reaction of a disuifide bond(s) in polypeptide, is directly exhibited. In addition, the present invention can be applied to a case in which the PDI activity exerts influence on proteins relating to gene expression, polypeptide folding or transport, thereby indirectly improving the productivity of PDI. According to the sembodiment of the present invention, the foreign gene is a gene coding for numer serum albuming

The term "polypeptide" as used herein means a short- or longchain peptide and a protein.

Examples of hosts include prokaryotes such as E. coli, Bacillus subtills, etc and sukaryotes such as yeast, animal cells, etc. preferred is a host cell capable of secreting a mature polypeptide through post-translational modification and processing, more preferably eukaryotes, and most preferably yeast.

The present invention also provides a process for producing a polypeptide, which comprises the following steps of:

co-expressing a human PDI game and a foreign game coding for the polypeptide to be produced, in the above-described transformant so as to produce the polypeptide; and

recovering the polypeptide

then HSA and ONI were co-expressed in an HSA-producing yeast

strain transformed with a human FOI expression glasmid in an appropriate medium according to the embodiment of the present invention, a secretion level of HSA was practically increased up to

bout 60% in average in comparison with the case of a non-ransformed HSA-producing yeast strain (pAH/HIS23) (Fig. 8).

Although we do not intend the present invention to restrict by theory, the increase in the secretion level of RSA by co-expression can be explained as follows:

HSA is a protein containing 17 disulfide bonds. It is known also that formation of its higher-order structure is enhanced in the presence of a stoichiometric amount of PDI in in vitro refolding apperiments of a denatured protein.

polyacrylamide gel electrophoresis, it was directed as a single band to-express a foreign PDI cDMA together with MSA. This indicate that of enhances the formation of a normal dispifile bond(s) in the MSA a water-soluble presence of a reducing agent, while detected as discontinuous bands SDS-polyacrylamide intracellular HSA sample prepared from a yeast strain which can not having a larger molecular weight than normal HSA in the absence of decule, but some of the HSA molecules are also detectable within intracellular HSA molecules is based upon the incomplete formation an intramolecular disulfide bond(s). In a yeast strain allowed to co-express together with MSA, however, an intracellular HSA οť he yeast cell. When the intracelluler HSA was analyzed by SDSjel electrophoresed without a reducing agent when compared with ceducing agent, clearly showing a different behavior from that Ath the same mobility as that of a normal HSA molecule in the miscule and thereby assists the formation of the higher-order normal HSA. These results indicate that the presence of ŝ HSA is secreted from the yeast strain HIS23 nolecule was detected as a more narrow band on

structure of ESA molecule more efficiently. Accordingly, it is suggested that the co-expression of FOI reduces chances of Causing the association of RGA molecules having unstable structure and their decomposition by protesses in the bost Call, thereby increasing the secretion of RSA molecules.

molecules by their coexistence in endoplasmic reticulum, because the based upon the fusion of the PDI gene with the MSA prepromaguence expressed. These results suggest that PDI exerts influence not only directly on ASA molecules but also on the transcription level of the increased lavel of the secretion of human FDI out of the yeast cell production level is based on the direct influence of PDI on HSA However, it seems to be reasonable that the increment of of enough amount to enhance the in vitro refolding of a denatured HSA is localized in the endoplasmic reticulum of the yeast cells in an also higher than ASA. These results, therefore, indicate that PDI which plays a role in the intracellular transport into endoplasmic cells are compared reticulum through a cransmenbrane process. In addition, when the than HSA, and the layel of human PDI detectable within the cells with each other, PDI is secreted in several times larger amounts expression by means of northern blotting, increase in the amount MRNA can be found in the former cells in which the PDI gene When the amount of HSA mRNA in the HIS23 in which PDI was expressed is compared with the amount in a control without coincrease in the amount of secreted 83% has a cornelation to which also supports the direct effect of PDI on amounts of HSA and Pul secreted from the HIS23 gene. RSA HSA



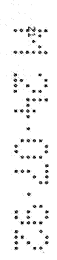
order structure of HSA. based on the direct influence of PDI on the formation of the higherexpression of POI on an increment of the amount of secreted MSA is highly amplified co-expression of PDI in the same host cell. proteins in which the formation of a disuifide bond(s) contributes improving effect can also be expected in other general secretory to the formation and stability of their higher-order structures, Thus, it is highly possible that the effect of the In consequence, such a similar secretion-Ċ

illustrate the present invention. following non-limited examples will be provided to further

[Examples]

Cloning of human PDI (protein disulfide isomerase) cDNA

0.28 maltose. After further adding 5 μl of 1 κ MgCl2 solution N, Amersham). With the phage-attached side upward, the filter was Bacto-trypton, 1% MaCl and 0.5% yeast extract) supplemented with the E. coli cells with the phage particles. The resulting cells thereto, the mixture was incubated at 37°C for 10 minutes to infect Y1090 which has been precultured overnight at 37°C in LB medium (1% The phage particles obtained were transferred onto a filter was incubated overnight at 37°C so as to grow the phage particles. were added to 50 ml of an LB top agar medium (LB medium, 10 mM MgCl2 (23 (Clontech) were mixed with 500 μ l of a culture of E. coli strain cm \times 23 cm). After solidifying the top agar medium, the plate 0.7% agarose), and then mixed and inoculated on a 18 agar plate About 100,000 clones of human liver Agtil cDNA library



Using the filter thus obtained, a screening of human PDI cONA was soaked in a neutral solution (I M Tris-RC1 (pH 7.5) and 1.5 M MaC1). then for further 1 minute on the same filter paper which has been soaked in an alkaline solution (0.5 N NaOH and 0.15 M NaCi), and put for 1 minute on a 3MM filter paper (Whatman) which has been carried out according to the following procedure: Thereafter, the filter was washed with 2 imes SSC solution (20 imes SSC imesM MaCl + 0.3 M trisodium citrate), air-dried and then exposed to ray for 2 minutes so as to fix the phage DNA on the fliter

proline-4-hydroxylase (the same protein as PDI) cDNA (Pihlajaniemi, automatic DNA synthesizer (Model 380B, Applied Biosystems). the complementary chain of the 243-282 base sequence of human TGGCGICCACCTTGGCCAACCTGATCTCGGBAACCTTCTGC-3') which corresponds et al., EMBO J., vol.5, p.543, 1987) was synthesized using an As a probe to be used, a 40 mer oligomer DNA (3)

fm/bd Ficel 400 + 2% polyvinyl pyrrelidene), 1 M NaCl, 50 mM Tris-HCl (pH further scaked in a hybridization solution (prepared by solution (100 x Denhardt solution = 2% bovine serum albumin + 2% the prehybridization solution which consists of 5×0 enhards Amersham) and 12 units of T4 polymuclectide kinese (Takara Shuzo, MgCl2, 5 mM dithiothzeitol, 100 µCi (y-32p] ATP (~ 3000 Ci/mmol, minutes in 50 µl of 50 mM Tris-HC1 (pH 7.5) buffer containing 10 phosphorylation, by incubating 20 pmoles of the DNA at 37°C e H The 5'-end of the synthesized DNA was labeled by 24 an ultrasonic-treated salmon sperm DNA. mM EDTA (pH 8.0), 0.1% sodium dodecyl sarcosinate and The filter obtained above was soaked at 37°C for I hour in The filter was 3



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positive phages from the primary screening were further subjected to was further subjected to a third screening in order to isolate a second screening under the same conditions as those of the primary cpm/mj of The resulting sections in 1 ml of SM buffer (100 mM NaCl, 10 mM MgCl2, 50 mM Trisminutes, followed by its exposure to an X-ray film (XAR-5, Kodak) (pH 7.5) and 0.01% gelatin), and left overolight at 4°C so as aforesaid particles with 2 x SSC + 0.1% sodium dodecyl sarcosinate solution at $42^{\circ}\mathrm{C}$ and This plate by cutting it out as gel sections, soaked each of the gel development of the film, 8 positive recover the phage from the gel into the solution. When the 8 filter was washed with 2 x SSC solution at room temperature $^{\mathrm{supplementing}}$ the prehybridization solution with about $^{\mathrm{10}6}$ screening, only one of them remained as a positive clone. Phage corresponding to those signals were recovered from the 37°C. hours at signals were detected by the primary screening. u) the aforementioned labeled DWA) for it as a homogeneous positive clone overnight at -80°C. After

20 units of EcoRI (Wippon Gene, Japan). By 0.8% HE. inally by the method of Leder et al. (Leder, P., Tiemeir, D. and Ç. it was found of about phage DNA (1/5 vol) was digested at 37°C for 1 hour in 50 µl of prepared MaCl, 6 mM MgCll, 6 mM mercaptoethanol, 0.1% gelatin, 20 µg/ml digestion salution consisting of 100 mM Tris-HCl (pH 7.5), 100 The insert DNA was separated and purified using glass A phage DNA was prepared from the positive clone obtained fragment thus agarose gel electrophoresis of the resulting digest, that this positive clone contains an Insert DWA Enquist, L., Science, vol.186, p.175, 1977). ribonuclease A and pb. 150

IB medium supplemented with 25 µg/ml of ampicillin, and a plasmid resulting mixture was then incubated at 16°C for 15 hours to obtain 1979). 10 µg of the plasmid DNA was digested at 37°C for 1 hour in 걸 ul of the digestion solution consisting of 100 mM Tris-RCI (pR screening. out by the Mandel's method (Mandel, added to the mixture of Liquid A 20 µl and Liquid was purified from the cultured ceils by alkaline lysis method about 100 ng of puci9 vector which has been digested mercaptoethanol, 0.1% gelatin was recovered by glass powder recovered DNA 100 extracted with phenol, concentrated by ethanol precipitation and hen subjected to 0.8% agarose gel electrophoresis. Thereafter, (Birmbolm, B.C. and Doly, J., Muciaic Acids Res., vol.7, p.1513). together. Using 10 µl of this reaction mixture, transformation recombinant plasmid in which both DWA fragments were linked ul from the DNA ilgation kit (Takara Shuro, Japan), and the 37°C in The digest was technique, for use as a probe in the following PDI cDMA transformant thus obtained was cultured overnight at p.154, 1970) powder (Gene Clean^{2M}, Bio-101). About 20 ng of the and 100 units of EcoRI (Nippon Gene, Japan). and Higa, A., J. Mol. Biol., vol.53, 7.5), 100 mM NaCl, 6 mM MgCl2, 6 mM insert DNA fragment of about 150 bp coll strain TG1 was carried and

In order to obtain a clone which contains the full length human PDI cDNA, screenings were carried out egain from about 50,000 clones of human liver Agtll cDNA library and about 50,000 clones of human placenta Agtll cDNA library were fixed were prepared in the same manner as described in the foregoing. In this instance, about 100



0 restriction maps of these clones with that reported by Pihlajaniemi placenta-originated estimated that the full length human PDI cDNA desired is covered placenta cDNA library. library were found to overlap one another. from the liver cDNA library and 2 clones from the placenta separately subcloned into an EcoRI site of plasmid vector pUC19 The EcoRI insert DNA fragments of the obtained 7 clones were third screenings, 4 positive closes were isolated from the liver CDNA library, and 3 positive closes from the placenta cDNA library at -80°C. the labeled DNA) for 15 hours at as a probe in the screening. After soaking the above two filters translation kit (Amersham), and the labeled cDNA fragment was used Bu found from the liver cDNA library, and 5 positive signals from the followed by their exposure to X-ray films (XAR-5, Kodak) overnight SEC + 0.1% sodium dodecyl sarcosinate solution at 65°C for 1 hour, washed with 2 imes SSC solution at room temperature and then with 0.5 supplementing the prehybridization solution with about 106 cpm/ml of filters were labeled using $\{\alpha^{-32}P\}$ dCTP (>400 Ci/mnol, Amersham) and a of the aforementioned 150 bp human PDI cDNA fragment was isotopethe inserts aforementioned prehybridization solution for 1 hour at 60°C, DNA base sequences of the two clones were determined using manner liver-originated clones (pHDFI16) and After development of the films, 5 positive signals were further soaked in a hybridization solution (prepared of the 7 clones. as described above in order to make restriction maps clones (pHDPIp4) based upon the comparison By subjecting these clones to second and 60°C. As the results, 4 clones obtained The resulting filters were In addition, it one Ö, nick



M13 SEQUENCING MIT (Toyobo, Japan.), M13 Sequencing Kit (Takara Shuzo) and an automatic DNA sequencer (370A, Applied Biosystems). Comparison of the thus determined sequences with the data reported by Pihlajaniemi et al. confirmed that the full length human PDI cDNA which consists of 2454 base pairs is encoded by these two clones (SEQ ID MO:1).

Construction of plasmid for human PDI expression in yeast

A plasmid for use in the expression of human PDI in yeast was constructed by the following procesure, using the above two clones, pHPDI16 and pHPDI94, which encode human PDI cDNA (Figs. 1A, 1B and 1C):

1.3 kb which corresponds to a 3'-end Separately from this, about 1 µg of pHPDIp4 DNA was digested at 37°C corresponds to a 5'-end EcoRI-Patl fragment of the PDI cDNA. 0.1% gelatin, Tris-HOl (pH 7.5), 100 mM NaCl, 6 mM MgClz, 10 units of PstI (Nippon Gene). The resulting digest was subjected mercaptoethanol, 0.1% gelatin, 10 units of EcoRI (Nippon consisting of 10 mM Tris-HCl (pR 7.5), 100 mM NaCl, 6 mM was digested at 37°C for 1 hour in 20 µl of the digestion solution 0.8% agarose gel electrophoresis and then to the glass powder 1 hour in About 1 mg of purblic DNA prepared by the alkaline lysis method to separate and purify a DNA fragment of about 490 bp which Gene). 10 units of PstI (Nippon Gene) and 10 units of BamHI $20~\mu\text{l}$ of the digestion solution consisting of 10 mM The resulting digest was treated to separate and purify a PstI-BamHI fragment 6 mM mercaptoethanol, XXC fragment in the same manner Ġ, MgCl₂, 6 mM of the PDI



mpicillin) of 90 mm in diameter. White colonies grown on the plate Ġ supplemented Isaline lysis method, and the DNAs were analyzed using restriction ere picked up, plasmid DMAs were prepared from the colonies by tha by incubating for each) shuzo). With 10 µl of the reaction mixture obtained, a competent of plasmid vector pUC 119 which has Mg/ml of 5-bromp-4-chloro-3-indoly1-8-0-galactopyranoside was cultured overnight at 37°C strain MV1190 cell was transformed by the calcium chloride \exists ug/m1 (iguid A and 5 µl of Liquid B of the DWA ligation kit (Takara mixture of 25 nzymes, thereby selecting a transformant which carries a 50 ng an X-Cal place (18 medium containing 1.5% agar further $\mu g/ml$ of isopropyl- β -0-thiogalactopyranoside and 25 The thus obtained plasmid was named phPDIEB BanHI, The thus recovered two DNA fragments (about the form with EcoRI and for 15 hours in cechnique. The transformed cell 20 29 digested in a linear samples at 16°C vere ligated with about lesmid. with 50

Calcium rypton, 0.5% NaCl and 1% Bacto-Yeast Extract) supplemented with 150 into the boundary region between the PDI signal sequence and the PDI Using the plasmid phPDIEB, a Nael cleavage site was introduced When turbidity (OD660) g/ml of ampicillin. One ml of the pre-culture was inoculated into hloride rechnique. A single colony of the resultant transformant coli strain equence itself on the cDMA by the method of Kunkel (Kunkel, T.A., Bacto-2 DNA as pre-cultured overnight at $37^{\circ}\mathrm{C}$ in 2 x YT medium (1.6% 0 ml of the 2 x YT medium supplemented with 150 $\mu g/ml$ of competent cell was transformed with the phPDIES vol.82, p.488, 1985). mpicillin, followed by its culture at 37°C. Natl. Acad. Sci., USA,



(m.o.i. * 2) was added aquecus layer was then mixed with 1/10 volumes of 3 M sodium acetate µg/m1, mixed with an equal volume of neutral phenol with stirring, and then W. centrifuged to recover an aqueous layer. To the layer was added an The mixture was further room temperature. TE buffer (pH 8.0) consisting of 10 nM Tris-HCl and 1 mM EDTA, and 2.5 volumes of ethanol. After stirring, the mixture was left The DNA was washed with 70% ethanol, a carried out by incubating supernatant recovered was mixed with 1/5 volumes of a solution suspension was added kanamycin to a final concentration of 70 precipitate obtained by centrifugation was dissolved in 5 for 30 minutes at -80°C, followed by centrifugation in order dried under a reduced pressure and then dissolved in 100 µl by culturing at 37% for 20 hours with shaking. obtained culture was subjected to centrifugation, and the glycol #6000. subjected to centrifugation to recover an aquedus layer. 0 minutes at without shaking. medium reached around 0.3, M13K07 phage containing 2.5 M NaCl and 20% polyethylene equal volume of chloroform with stirring. stirring, the mixture was left for 15 and the infection was minutes recover DWA as precipitate. 9 Tor 37°C medium, U N buffer ollowed

Using the resulting phPDISB-originated single-stranded DNA containing dU, a desired mutation, i.e., introduction of a Nael site, was carried out in the following manner:



After adding 1 medium (2% Bacto-trypton, 0.5% yeast extract, 10 13 Takara Shuzo), 60 of the phosphorylated synthetic oligonucleotide solution obtained mW dithiothreitol, 1 mM solution which consists of 100 mM Tris-ACl (pH 8.0), 10 mM MgCi2, 7 BMH71-18mutS obtained reaction was terminated by adding 3 μl of 0.2 M EDTA (pH 8.0) and Shuzo) and 1 unit of T4 DNA polymerase (Mutan TM-K, Takara Shuzo), elongation solution (Site-directed mutagenesis system MutanTM.N., 37°C for 15 minutes. Thereafter, a complementary chain synthesis above, and the mixture was left at 65°C for 15 minutes and then to deactivate the T4 polymucleotide kinase. Separately from this, heating the mixture at 65°C for 5 minutes. 3 $\mu 1$ of the DNA solution and by incubating the resulting mixture at 25°C for 2 hours. The was carried out by mixing the reaction mixture with 25 μi of a chain system MutanTM-K, Takara Shuzo) were mixed with sterile water to (Takara Shuzo), followed by heating at 70°C for 10 minutes in order and 1 µ1 of an annealing buffer (Size-directed mutagenesis pmol of the above-described phpDIRE-originated single-stranded 10 mM MgSO4, 10 mM MgCi2 and 20 mM giucose), and the mixture volume of 10 µ1. thereto, ice bath. in an ice bath, was mixed with 30 µl of a suspension of E. coli strain at 37°C competent cell, and the cell suspension was left for 吕 and the mixture To of 2 x units of E. coli DNA ligase (Mutan TM-K, Takara for 1 hour. the cell for 45 One #1 ATP and 10 units of T4 polynucleotide kinase YT medium containing 150 µg/ml suspension seconds at of this solution was mixed with 1 µ1 10 µ1 10 11 11 11 of M13K07 phage was then for 42°C 30 minutes at and then for I minute mM WaCL, added 300 µl of soc 8.5% further 3



ampicillin and 70 µg/ml of kanamycin to the mixture, the mixture was shaken at 37°C for 20 hours. The resulting culture was centrifuged to recover 20 µl of supernatant which was subsequently mixed with 80 µl of a culture of £. coli strain MV1190. After incubation at 37°C for 10 minutes, the resulting mixture was inoculated onto a LB plate supplemented with 150 µg/ml of ampicillin and cultured overnight at 37°C. Among colonies grown on the place, a transformant carrying a Nasi site-introduced plasmid was identified by DNA-sequencing using w13 sequencing kiT (Toyobo). This plasmid was named phPDINae.

2 μg of the phPDINae DNA prepared by the aikaline lysis method was digested at 37°C for 4 hours in 30 μl of the digestion solution consisting of 10 mM Tris-RCl (pH 8.0), 20 mM NaCl, 7 mM MgClz, 7 units of MaeI (Nippon Gene) and 10 units of HindIII (Takara Shuzo). The resulting digest was subjected to 0.8% agarose gel electrophoresis and then treated by the glass powder technique so a co-separate and purify a DNA fragment of about 1.7 kb.

Shuzo), and Tris-HCl (pH 7.5), utilized in yeast was constructed in the following manner (Fig. 1A): deactivate the enzyme. human serum albumin prepro-sequence and is composed of codons often (Nippon Sene), followed by heating at 70°C for 5 minutes in order plasmid, pUC1198ig, containing a DNA fragment which encodes water and 1 unit 0 £ Ë the mixture was incubated of plasmid vector pUC119 DNA was digested at 37°C for 1 of the digestion solution which consists of 100 10 mM MgCl₂, 50 mM NaCl and 12 To the reaction mixture was added 38 of bacterial alkaline phosphatase DI FT 37°C for 1 hour, followed units of E Mar.

by phenol extraction

and ethanol precipitation to

recover DNA.

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DNA was then incubated overnight at 16°C in 30 µl of the ilgation solution which consisted of 66 mM Tris-HCl (pH 7.5), 6.6 mM MgCl2, 10 mM dithiothreitol, 0.1 mM ATP and 300 units of T4 DNA ilgase (Takara Shuzo), together with an equal molar amount of a Khoi linker containing a Khoi site and consisting of the following sequence: 5°-AATTCTCGAG

GAGCTCTTAA-5'

Using 10 µl of this solution, transformation of E. coll JM107 competent cell was carried cut by the calcium chioride method. The transformed cell was cultured overnight at 37°C on a LB plate supplemented with 50 µg/ml of ampicillin. Plasmid DNAs were prepared by the alkaline lysis method from the colonies on the plate and analyzed using restriction enzymes. In this way, a plasmid DNA molecule in which the XhoI linker has been inserted into pUCl19 EcoRI site was selected.

The following four types of oligonuclectides:

- (1) S'-TCGAGAATTCATGAAGTGGGTTACCTTCATCTTGTTGTT-3';
- (2) 5'-ARCABGRACARCARGRAGARGRAGGTRACCCACTTCATGARTTC-3';
- (3) 5'-crrgirorcrrccracrcracecrrrrcagaaGGCCTG-3'; and
- (4) 5'-GAITCCAGGCCTTCTGAAAACACCTCTAGAGTAAGCAGAAGAG-3'

were synthesized using an automatic DNA synthesizer (3808, Applied Siosystems).

sach 5'-end of these oligonucleotides was phosphorylated by incubating about 30 pmol of each sample at 37°C for 1 hour in the solution consisting of 50 mM Tris-HCl (pH 7.6), 10 mM MgClg, 5 mM lithiothreitol, 0.2 mM ATP and 6 units of T4 polynucleotide kinase (Takara Shuzo). The oligonucleotide solutions obtained were

combined (100 µ1 in total volume) and annealed by leaving the combined solution for 5 minutes in a water bath of 100°C, followed by cooling down to room temperature. To the solution was then added 600 units of 74 DNA ligase (Takara Shuzo), and the mixture was left overnight at 16°C to ligate these fragments. The double-stranded DNA preparation thus obtained was subjected to phenol extraction in order to remove proteins, and then to ethanol precipitation to recover the DNA.

followed by phenol extraction and ethanol precipitation to recover a S The fragment obtained was incubated overnight at 16°C lasmid DWAs prepared from colonies on the plate were analyzed using WAS units of T4 DWA ligase (Taxara Shuzo), together with an equal molar amount of the double-stranded DNA fragment obtained by ligating the 30 µl of the ligation solution which consists of 66 mM Tris-HCl carried out by four oliganucleotides. Using 10 µl of the thus prepared solution, consisting of 100 mM Tris-HCl (pH 7.5), 10 mM MgCl2, 100 mM MaCl, units of Bamfi (Nippon Gene) and 12 units of Mhol (Takara Shuzo), overnight at 37°C on is medium containing 50 $\mu\text{g/ml}$ of ampicillin One kg of the above XhoI linker-introducing vector plasmid digested at 37°C for 1 hour in 20 µl of the digestion solution restriction enzymes so as to select a transformant containing PH 7.5), 6.6 mW MgCl2, 10 mW dithiothreitol, 0.1 mM ATP and The obtained plasmid was named transformation of E. coli JM107 competent cells was The transformed cells the calcium chloride method. desired recombinant plasmid. DMA fragment.



piate 3.2 kb DNA fragment (about 50 ng) from pUC1195ig at 16°C for 30 electrophoresis and then treated by the glass powder technique to HindIII (Takara Shuzo), subjected to 0.8% agarose gel digestion solution which consists of 10 mM Tris-HCl (pH 8.0), 100 method. 2 μg of the DNA was digested at 37°C for 4 hours in the plasmid mixture of Liquid A 30 µl + Liquid B 5 µl). Using 10 µl of minutes in the ligation kit solution of Takara Shuzo(Japan) (a separate and purify a DNA fragment of about 3.2 kb. The 1.7 kb DNA reaction mixture, transformation of E. coll HE101 competent cells fragment (about 50 ng) derived from phPDINae was reacted with side of the human serum albumin prepro-sequence (Fig. 2). The prepared by the alkaline lysis method from colonies grown on the obtained plasmid was named phPDILyl supplemented with 50 µg/ml of ampicillin. Plasmid DWAs were transformed cells were cultured overnight at 37°C on a LB plate (Takara Shuzo) was carried out by the calcium chloride method. The and analyzed using restriction enzymes to select a recombinant DNA was prepared from plasmid pUC119Sig by the alkaline lysis mm MgCl2, 8 units of Stul (Nippon Gene) and 10 units of in which the human PDI itself was linked to the downstream the

A human PDI expression plasmid was constructed in the following manner, such that the leader sequence modified type PDI can express under the control of a promoter of yeast alcohol dehydrogenase I

gene:
7 µl of phPDILyi DNA prepared by the alkaline lysis method was
digested at 37°C for 2 hours in 100 µl of the digestion solution
which consists of 100 mM Tris-HCl (pH 8.0), 100 mM NaCl, 7 mM MgCl;



stirring, the mixture was centrifuged to recover an aqueous layer. mixed with an equal volume of a phenol/chloroform mixture (a mixture and 40 units of EcoRI (Rippon Gene). The resulting solution was The phenol/chloroform extraction was repeated, and the aqueous layer and 2.5 volume of ethanol. The mixture was left for 2 hours at obtained was mixed with 1/10 volume of 3 % sodium acetate (pH 5.3) in 50 µl of Klenow buffer solution (Deletion Xit for Kilo-Sequence, with 70% ethanoi, dried under a reduced pressure and then dissolved 0 phenol/chloroform extraction, and the resulting aqueous layer was site. The thus prepared solution was subjected twice to incubated at 37°C for 45 minutes to blunt-end the EcoRI cleavage units of Klenow fragment (Takara Shuzo), and the mixture was Takara Shuzo). Thereafter, to the solution obtained was added 4 $40^{\circ}\mathrm{C}$ and then subjected to centrifugation. The pellet was washed mixed with 1/10 volume of 3 M sodium acetate (pH 5.3) and 2.5 volume and the resulting aqueous layer was mixed with 1/10 volume of 3 M mixture was then subjected twice to phenol/chloroform extraction, Klenow fragment (Takara Shuzo), and the mixture was incubated at Shuzo). Thereafter, to the solution obtained was added 4 units of of Klenow buffer solution (Deletion Kit for Klio-Sequence, Takara ethanol, dried under a reduced pressure and then dissolved in 50 Hl subjected to centrifugation. sodium acetate (pH 5.3) and 2.5 volume of ethanoi. left for 1 hour at -40°C and them centrifuged. saturated phenol with an equal volume of chioroform). ethanol. The mixture was left for I hour at -40°C and then 100 45 minutes to blunt the Booki cleavage site. The pellet was washed with 70% 13. (b) peliet was The mixture was The reaction



precipitate was washed with 70% Ç E Thereafter, the DNA solution obtained was subjected to 0.8% solution which consists of 10 mM Tris-HC ro, was added 4 units of of Klenow buffer solution (Deletion Kit for Kilo-Sequence, Takara mm MaC1, 7 mm MgC12 and 24 units of KhoI (Takara Shuzo). The purify a DNA fragment of about 1. acetate (pH 5.3) and 2.5 volume of ethanol. This mixture of 10 prepared by ethanol, dried under a reduced pressure and then dissolved in 50 nixture was then subjected twice to phenoi/chloroform extraction, for 2 hours in 100 µ1 ethanol, dried under a reduced pressure and then gel electrophoresis and then treated by the glass powder digestion solution which consists of 10 mM Tris-HCl (pH 8.0) 3 M sodium acetate (pR 5.3) and 2.5 volume of ethanol. aqueous layer separated was mixed with 1/10 volume of 3 Separately from this, 5 µl of pubs-abh-ash-a DNA (Japanese extraction, and the aqueous layer separated was mixed with 1/10 pellet The mixture was left for 2 hours at $-40^{\circ}\mathrm{C}$ and then centrifuged reaction mixture was them subjected twice to phenoi/chloroform 60 mM NaCl, 7 mM MgCl2 and 10 units of BamHI (Mippon was incubated removed was washed with 70% ethanol, dried under a reduced and then dissolved in 40 µl of the solution which consists The The DMA 37°C for 45 minutes to blunt the KhoI cleavage site. Patent Application Laying-Open (ROKAI) No. 2-117384) Klenow fragment (Takara Shuzo), and the mixture left for 1 hour at -40°C before centrifugation. obtained alkaline lysis method was digested at 37°C recover DNA as precipitate. The DWA solution to separate and to the of the Shuro). Thereafter, 3 70% 12 40 (1) (0) 80 washed with rolume of issolved PH 8.0), cechnique



µ1 + Liquid B 6 µ1) in order to ligate the two DNA fragments. Using pellet alkaline phosphatase from E. coli strain C75 (Takara Shuzo), and ona ligation kit solution of Takara Shuzo (a mixture of Liquid A 30 mixture was incubated at 50°C for 1 hour in order to carry out 5'-end dephosphorylation of the restriction enzyme-formed cleavage To the reaction mixture was added 1/10 volume of 3 M sodium of the TS buffer. Thereafter, the prepared DMA solution was mixture obtained ã , 1 Q prepared by the alkaline lysis method from colonies grown on the (Nippon Gene). The solution obtained was incubated at $37^{\circ}\mathrm{C}$ for separated was dried under a reduced pressure and then dissolved modified type PDI sequence was linked to the downstream side of plate and analyzed using restriction enzymes in order to select IIIS-EC1 (pH 8.0), 60 mM MaCi, 7 mM MgCl2 and 10 units of BamHI subjected to 0.8% agarose gel electrophoresis and then treated ragment of about 8 kb. The thus obtained phPDILyl-originated prepared DNA solution, transformation of E. coll to separate and purify a DNA left for 1 hour at -40°C before centrifugation. The DNA minutes to digest the DNA. To the reaction mixture was then supplemented with 50 µg/mi of ampiciliin. Plasmid DNAs were transformant carrying a plasmid in which the leader sequence kb DNA fragment (about 50 ng) and pJDB-ADH-RSA-A-originated of 2 M Tris-HCl (pH 8.0), 110 µl of sterile water and fragment (about 50 ng) were incubated at 16°C for 2.5 hours T.E 0600 was carried out by the calcium chloride technique. transformed cells were cultured overnight at 37°C on a 322 scetate (px 5.3) and 2.5 volume of ethanol. powder glass technique so as of the -1 펿



alcohol dehydrogenase I promoter. The constructed PDI expression plasmid was named pAHhPDILyl. As the results of the plasmid construction, the N-terminal amino acid of the mature PDI protein was changed from Asp to Gly.

A control plasmid for use in experiments of the human PDI expression was constructed in the following manner:

pressure and dissolved in 20 µl of the TE DNA solution was subjected to 0.8% agarose gei electrophoresis and mixed with 1/10 vol of 3 M sodium acetate (pH 5.3) and 2.5 phenoi/chloroform extraction, and the aqueous layer separated mixture obtained was then left for 2 hours at ~40°C and then 1/10 vol of 3 M sodium acetate (pH 5.3) and 2.5 vol of ethanol. phenol/chloroform extraction, and to the aqueous layer was added solution which consists of 10 mM Tris-HCl, 100 mM NaCl, 7 mM MgCl2. method was digested at 37°C for 2 hours in 100 μl of the digestion cleavage sites. The reaction mixture was then subjected twice incubated at 37°C for 45 minutes to blunt the XhoI and BamMI Takara Shuzo). Thereafter, to the obtained solution was added a 4 in 50 µl of Klenow buffer solution (Deletion Kit for Kilo-Sequence with 70% ethanol, dried under a reduced pressure and then dissolved centrifuged to recover DNA as pellet. The DNA pellet was washed Gene). The reaction mixture obtained was subjected twice to 24 units of Whol (Takara Shuzod) and 29 units of BamHI (Nippon of Klenow fragment (Takara Shuzo), and the mixture was The mixture was left for 1 hour at -40°C of pJDB-ADH-HSA-A DNA prepared by the alkaline lysis The DNA pellet removed was dried under a reduced buffer. Thereafter, the before The



then treated by the glass powder technique to separate and purify a DNA fragment of about 8 kb. The thus obtained DNA fragment (about 50 ng) was mixed with the mixture of Liquid A 30 µl + Liquid B 5 µl from the DNA ligation kit (Takara Shuzo), and incubated overnight at 16°C so as to cyclize it by self-ligation. Using 10 µl of the prepared DNA solution, transformation of E. coli 101 competent cells from transformed cells were cultured overnight at 37°C on a LB plate supplemented with 50 µl/ml of ampicillin. Plasmid DNAs were prepared by the alkaline lysis technique from the colonies grown on the plate, and analyzed using restriction enzymes in order to select a desired control plasmid. The constructed plasmid was named pAH.

Expression of human EDI in yeast

Using the human PDI expression plasmid pAHhPDILYi constructed above, an expression of human PDI in yeast was carried out in the

A single colony of yeast strain AH22 obtained by culturing it on a YPD plate (2% Bacto-pepton, 1% yeast extract, 2% glucose and 1.5% agar) was inoculated into 5 ml of a YPD medium (2% Bacto-pepton, 1% yeast extract and 2% glucose) and cultured at 30°C for 24 hours with shaking. This pre-culture (0.9 ml) was then inoculated into 45 ml of the YPD medium and cultured at 30°C with shaking. When turbidity at ODeon reached about 0.5, the main culture was subjected to a low speed centrifugation to recover yeast cells as precipitate. The cells removed were suspended in 3 ml of 0.2 M Lisco, and the cell suspension (1 ml) was centrifuged to recover the

inoculation of its suspension onto a SD(-Leu) plate (SD(-Leu) medium mg/1 of threonine and 375 mg/l of serine (amino acids from Wako Pure cultured at 30°C for 24 hours with shaking. 1,5 ml of the resulting 5-days culture was inoculated into 5 ml of the SD (-Leu) medium and main culture was centrifuged to recover 500 µl of supermanant which Bromophenol Blue). After boiling for 5 minutes, the treated sample To the cells were subsequently added 46 µl of 50% PEG #4000 The pellet was resuspended (0.67% Bacto-nitrogen base, 2% glucose, 20 mg/l of adenine, 20 mg/l Chemical Industries, Japan)) + 1.5% agar). A transforment from the dissolved in 10 µl of a sample buffer for SDS-PAGE (125 mM Tris-HCl iscleucine, 30 mg/l of lysine, 50 mg/l of phenylalanine, 100 mg/l oultured at 30°C for 2 days with shaking. 100 µl of the obtained aspartic acid, 100 mg/l of glutamic acid, 150 mg/l of valine, 200 was subsequently mixed with the equal volume of ethanol and then arginine, 20 mg/l of methionine, 30 mg/l of tyrosine, 30 mg/l of 41 of Liscw and 10 µl of a pashebily 1 DNA solution (27 µg as pre-culture was then inoculated into 5 ml of the YPD medium and to recover products secreted out of the yeast cells as a pellet (pH 6.3), 4% SDS, 20% glycarol, 10% \$-mercaptoethanol and 0.01% The pellet was pipetting, the mixture was left oversight at 30°C, followed by suspension in 1 ml of sterile water. The suspension was then for 1 hour in an ice bath. The mixture was centrifuged After mixing them by uracil, 20 mg/l of tryptophan, 20 mg/l of histidine, 20 in 100 µl of sterile water and cultured at 30°C after the which was then dried under a reduced pressure. centrifuged to recover cells as pelfer. prepared by the alkaline lysis method.



ဌ out Blue, 10% acetic acid reight, 94,000), bovine serum albumin (67,000), ovalbumin (43,000). subjected to electrophoresis on SDS/PAGS Plate 10/20 (Dailchi except that panhPDILy! was replaced by the aforementioned control plasmid pAH, was run at the same time during the electrophoresis. expressed and secreted. Next, a large-scale culture was carried product. as standard molecular weight markers, phosphorylase b (molecular carbonic anhydrase (30,000), soybean trypsin inhibitor (20,000) in the following manner in order to examine chemical properties mature PDI protein, it was assumed that a desired human PDI was and 40% methanol) and then soaked in a decoloring solution (10% this instance, a control sample obtained by the same procedure, expression product having a molecular weight of about 55 K was found. Since this molecular weight coincided with that of the The resulting gel was stained with a x-lactalbumin (14,000) were used (Fig. 3). As the results, scetic acid and 40% methanol) to visualize an expressed staining solution (0.15% Coomassie Brilliant expressed and secreted protein: Japan) Kagaku Yakuhin,

A single colony of the pARhPDILy1-carrying yeast strain AM22 was incculated into 80 ml of the SD (-Leu) medium and cultured at 30°C for 2 days with shaking. The obtained pre-culture was then inoculated into 4 liters of a YPD-phosphate medium (YPD medium, 6 g/l of Na2HPO4, and 3 g/l of KH2PO4, pH 7.0) and cultured at 30°C for 24 hours with shaking. The resulting main culture was centrifuged to removed the supernatant which was used for the purification of the secreted expression product.



Isolation of recombinant human FDI from the culture and its

characterization

yeast was concentrated to 1/40 (final volume, 100 ml) using a elution was carried out in 10 mm borate-10 mm xCl buffer (pH 8.0) hydrophobic column chromatography so as to isolate human PDI. The weight, 30,000 cut-off), and then subjected to TSK-gel Phenyl-SPW Millipore-Millitan ultrafiltration apparatus (nominal molecular ammonium sulfate over 125 minutes. containing 0.05% NaNg, with a linear gradient from 0.85 M to 0 M removed markedly efficiently by the chromatography. the hydrophobic column chromatography without a loss of in the figures, the human PDI was purified almost homogeneously by electrophoresis of the isolated human PDI is illustrated. As shown result is shown in Fig. 4. In culture (4 liters) obtained by culturing the recombinant Any UV-absorbing substance in the YPD medium could Fig. 5 the result of SDS-The flow rate was 2 ml/min.

PDI ASSAY

pDI assay was carried out by measuring its effect to enhance the refolding of scrambled ribonuclease A (RNase A) which has been prepared by reduction, denaturation and re-oxidation steps. Refolding degree of the scrambled RNase A was determined by measuring a degree of the restoration of its enzyme activity. The following describes the assay procedure illustratively: Preparation of scrambled RNase A:

120 mg of RNase A was dissolved in 3 ml of 0.1 M Tris-HCl buffer (pH 8.5) containing 6 M guanidine hydrochloride and 0.15 h



dithiothreitoi, and then reduced under nitrogen atmosphere at room temperature for 15 hours. The reduced product was applied to a Sephadex G-25 column (15 mms x 38 cm) equilibrated with 0.01 N HCl, thereby removing the reducing agent. To the desalting product was added guanddine hydrochloride to a final concentration of 6 M. After adjusting its pH value to 9.0 with Tris, the mixture was subjected to an exchange reaction of a S-S bond(s) in the dark at 4°C for 14 days. The thus prepared sample was stored at -80°C for use as the scrambled RNase A.

PDI assay:

my phosphate buffer (pH 7.5) mixed with 20 Hl of an enzyme sample, phosphate buffer (pH 7.5) in which any dissolving air was replaced with mitrogen gas. Separately from this, 1.945 mi of degassed 50 mM Tris-HCl buffer (pH followed by the enzymatic reaction at 30°C for 15.5 minutes. stirring, a temperature of the resulting mixture is maintained mM EDIA; adjusted to such a concentration that absorbance at 280 nm 7.5) containing 5 mM and, with stirring, a change in the absorbance of scrambled RNase A solution is added to becomes solution (dissolved in 10 mM Tris-RCl buffer, pH 7.5, containing 1 added 50 µl of the scrambled RNase solution prepared above, the mixture is left for 5.5 minutes at 30°C. 13 treatment. µ1 of 1 M dithiothreital is added to 20 ml 80) is placed into a quartz cell (1cm x 1cm). this instance, absorbance at 260 nm should not be changed Thereafter, 5 µl of the dithiothreital-treated 10 µl of this solution is added to 420 µl of 55 MgCl2, 25 mM KC1 and 50 µ1 of a yeast RNA the buffer in the quartz cell To this solution the reaction Ç, បា បា

mixture at 260 nm are measured over 2 minutes at 0.2 minute intervals. The PDI activity is calculated from an initial velocity of the changing rate of absorbance at 250mm.

Transformation of yeast strain RIS23 with numan PDI expression plasmid pakarplist

Using the aforementioned human PDI expression plasmid pawheDily1, transformation of the MSA-producing yeast strain MIS23 (Japanese Patent Application No. 2-57885 filed by the present applicant, Bikoken-Kin-Ki No. 11351 (FERM P-1138)) was carried out in the following manner:

at 30°C with shaking. When turbidity at OD600 reached about main culture was subjected to a low speed centrifugation to place (2% Bacto-trypton, 1% Bacto-DNA) prepared by the alkaline lysis method (Birnboim, H.C. and Doly, of a YPD medium (2% Bacto-trypton, 1% yeast extract and 2% glucose) To the pellet were added 46 µl 23 resulting centrifuged to of 50% polyethylene glycol #4000, 10 µl of Lisch and 10 µl of the obtained pre-culture was inoculated into 50 ml of the YPD medium J., Nucleic Acids Res., vol.7, p.1513, 1979). After mixing them yeast extracts, 2% glucose and 1.5% agar) was inoculated into 5 40 human PDI expression plasmid pathrPDILyl DNA solution (about 20 of the A single colony of the HSA-expressing yeast strain HIS23 \exists recover cells as pellet. The pellet was suspended in 100 One mi pipetting, the mixture was left overnight at 30°C. and and cultured at 30°C for 24 hours with shaking. mixture was suspended in 1 ml of sterile water redover the yeast calls as pallet. a YPD ទ obtained by culturing it 0.5, the

acid, 4 43, (0.67% Bacto-nitrogen base, mg/l of valine, 200 mg/l of threonine and 375 mg/l of serine (amino acids from Wako Pure Chemical Industries)) + 1.5% agar). rransformant was obtained as a colony grown on the plate on the tryptophan, 20 mg/l of arginine, 20 mg/l of methionine, 30 mg/l phenylalanine, 100 mg/l of aspartic acid, 100 mg/l of glutamic Q. 30°C after inoculating the cell mg/l of ispleacine, 30 mg/l of lysine, 50 mg/l mg/1 uracil, 20 37 20 mg/l suspension onto a SD (-His, -Leu) plate 2% glucose, 20 mg/l of adenine, at starile water and cultured after the culture tyrosine, 30 150

Expression of PDI in the obtained transformant pathyphily!/HIS23) was examined in the following manner:

In this instance, a transformant (pAH/HIS23) obtained using the 200 ul of the pre-culture under a reduced pressure using an evaporator. The precipitate Blue and 20% glycerol). After boiling for 5 minutes, the sample was then inoculated into 5 ml of the YPD medium and cultured at 30% the resulting main culture was Tris-HCl (pH 5.8), 2% SDS, 5% B-mercaptoethanol, 0.005% Bromophenol prepared by removing the PDI cDNA moiety supernatent which was subsequently from pAthPDILyl was used as a centrol. The single colony grown on -Tec) medium mixed with the equal volume of ethano! and then left for 1 hour secreted out of the yeast cells as a precipitate which was then was dissolved in 10 µl of a sample buffer for SDS~PAGE (62.5 mM an ice bath. The mixture was centrifuged to recover products plate was inoculated into 3 ml of the SD (-Mis, cultured at 30°C for 2 days with staking. 100 18.2 O.5 15 CH 24 hours with shaking, 1.5 plasmid pak which has been centrifuged to recover 500



subjected to electrophoresis on SDS-PAGE Plate 4/20-1010 (ex Dailchi Kagaku Yakuhin). The resulting gel was stained with a staining solution (0.15% Coomassie Brilliant Blue, 10% acetic acid and 40% methanol) and then soaked in a decoloring solution (10% acetic acid and 40% instance, phosphorylase b (molecular weight, 94,000 daltons), bovine serum albumin (67,000), ovalbumin (43,000), carbonic anhydrase (30,000), soybean trypsin inhibitor (20,000) and orlactalbumin (14,000) were used as standard molecular weight markers (Fig. 6).
As the results, the secretion of an expressed FDI having a molecular weight of about 55,000 daltons was detected in the yeast strain \$MIS23 transformed with pashpolly1.

Effect of human POI on the expression and secretion of HSA

Using the above-described co-expression system of HSA and PDI in a yeast cell, effects of human PDI on the expression and secretion of HSA were examined in the following manner:

Five single colonies were isolated from strain pAH/HIS23 which has been obtained by transforming the yeast strain HIS23 with the control plasmid pAH, and other 5 single colonies were isolated from strain pAHhPDILy1/HIS23 obtained by transforming the HIS23 with the human PDI expression plasmid pAHhPDILy1. Each of the thus isolated single colonies was inoculated into 5 ml of the SD (-His, -Leu) medium and cultured at 30°C for 24 hours. Each of the pre-cultures (100 µl) was inoculated into 5 ml of the YPD medium and cultured at 30°C for 24 hours with shaking. From the main cultures obtained, samples for SDS-PAGE were prepared in accordance with the



aforementioned procedure. Results of the SDS-PAGE are shown in Fig. 7. Using the gels subjected to the SDS-PAGE, the amount of secreted HSA from each strain was determined using a densitometer (IMAGE ANALYSIS SYSTEM, ex TEFCO, Japan) in order to examine effects of the co-expression of PDI on the secreted amount of expressed HSA (Fig. 8). As shown in the figure, the strains pAH/HISZ3 and pAHhPDILyl secreted HSA in average amounts of 0.93 mg/l and 1.50 mg/l, respectively. In other words, secretion of HSA was increased by about 50% in average due to the co-expression of PDI in the yeast strain HISZ3.

[Advantages of the Invention]

high-order structure thereof is incomplete because of the mistaken to enhance the activation of a protein in which the formation of a host mell by coupling both the espressions of the enzyme and the maans of generic engineering techniques, and it is possible to large amount of the protein economically. bond formation of a 5-5 bond(s), and thus as a means to prepare a isomerase gene. Accordingly, this process can be used as a means utilizing a fusion gene composed of a DNA fragment coding for human scale preparation of human PDI has been fast established by e ur aproduction advisous so so commissa unicompose a excitors invantion is the activation of an inactivated profess produced serum albumin pre-pro sequence and a human protein disulfide aseful polypeptide. according to the present invention, a process for the large-In addition, the PDI can A metr advantege of the be used as a reagent

SEQUENCE LISTING

SEQUENCE LENGIR: 2454 SEQ ID NO: 1

SEQUENCE TYPE: nucleic acid

STRANDEDNESS: double

TOPOLOGY: linear

to mRNA CDNA MOLECULE TYPE:

placenta Agtil cona libraries 800 INMEDIATE SOURCE: human liver

(Clontech)

SEQUENCE DESCRIPTION:

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Leu Arg

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53 900 TTC CTG CGG AMA AGC AAC Lys Ser 933 (5) [1] CIC GAC CAC G&G G&G GAG

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Phe

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8 300 THE CLE GAG GIG CEG 950 33.0 10 3.46 GCC CAC 808 810 808

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ec C Tyr P26 333 Leu Leu Val 23.5 27.3 3.1.35 3.1.35 Leu Aia Ala Gla

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24.9 T. GCC GCT Ala 2,73 GCC AAA 5. Tyr GCC CCT GAG TAT G.1 te 025 833 CIG Leu GCC CAC TGC AAG GCT なごせ 273 Cys (A) GLY 761 S 991 dir

297 GCC AAG GTG GAC COS ANG CIG ANG GCA GAR GGT TCC GAG ATC AGG TIG

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345 FEL 080 **61**3 Arg 260 80 GIG Val 088 Tyr Gly CAG TAC GIn CAG Gin A1.a CTG GCC Leu TCT GAC ASD in Si Glu GAG GAG G12 H 디 디 Ala

393 GAR 614 55 F AAG ပ္ပင္ပ Pro GGA SAC ACG GCT TCC Ser 95 Thr Als dsw Gly Asn AGG AAT Arg TTC Phe Fhe TIC ATC AAG 273 136 ri ri Pro 200

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633 AAA Lys COL 77. Phe Asa Ser Asp Val GIG GAC TIT 668 ATC ACT TCC AAC AGT Thr Ser Gly Ile 919 Ile Pro GAC ATA CCA 150 Asp

189 GAA 0.75 QS S GAT LLI Phe Lys > CIC TIT AAG AAG Ľýš Phe Leu V.2. GET GEC Val GAC AAA GAT GGG Gly Asp 5557 ren Asp CIC Gla

723 CTG GAC TIT GAA GGG GAG GTC ACC ANG GAG AAC CTG CGG AAC AAC 38C

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gay cag ana gov gyg ana gay san cig tan tacgcamage cagacecigg

1595

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asp Gln Lys Ala Val Lys Asp Glu Leu *

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2435 2454 CTACCGTGTT CSGAGTCTCG CTGCCTCCCT 1895 GACATTTTC 1955 2015 2075 ITCHACAGEA EGGECTGTGG CCTGTTCAAG GCAGNACCAC GACCCTTGAC TCCCGGGTGG 2315 SCARCTIGGEC ARGGATGETG GASCTGAATE AGACGETGAE AGTTETTCAG GEATTTETAT 2375 GOTROCCAS ACCCUTOSES SEUTSCACAC COAGOASOAS CSCACSCOTO CSAASCOTSC 1655 TTTCGARAG 1775 SEATCATGGO TOTTGCATTT TIGGGTAAAT GGAGAOTTCO GGATCOTGTO AGGGTGTODO 2135 caracerse aacacetes serseersee asceeress ecosseada secresseer 2195 ecrocrocro crategres crearrers coacrescer 2255 SANCCAICT CCAGGCAGCC CACOCTGGTG GGGCTYGTTT CCTGAAACCA YGATGTACTT 1835 probabite antigacae appeceara baractigar attitrocca cocarabaka SCCTOSCIT GAAGGAGGC STCGCGGAA ACCCAGGGAA CCTCTCAAA GTGACACTC receesar GONGGCACG GAGCGGGACT GGACATGGTC ACTCAGTACC GCCTGCAGTG TCGCCATGAC ACCCCTACAC ACCGTCCGTT CACCCCGTC TCTTCCTTCT GCTTTTCGGT COGOGGGAG SITICICCT INTITIGIARA ITCCGICIGI GGGAITITIA ARANCAGGS TATTIGITICS ACCITGECTA GGCCTCCTCG GAGAAGCTTG AGAGTGCTTG rrcaracar eagrerere recentree reasseases AAAAAAAA CCCGAATTC

SEQ ID NO: 2

SEQUENCE LENGTH: 1545

SQUENCE TYPE: nucleic acid

STRANDEDNESS: double

IOPOLOGY; linear

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A.A.G

AAG

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ACE GOT TOO COC AAG GAA

GGA GAC

Lit

ACC ATC AAG TTC TTC AGG

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83

Tyr Gly val Arg

Glu Ser Asp Leu Ala Gln Gin

GIn

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A.1.2

Ser Pro Lys Glu

Thr Ala

Ile Lys Phe Phe Arg Asn Gly Asp

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4.62

3

ACS SGC COS GCT SCC ACC ACC CTG CCT GAC SGC GCA GCT GCA GAG

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Six

Thr Gly Pro Ala Ala Thr Thr Leu Pro Asp Gly Ala Ala Ala

Ala Giy Arg Glu Ala Asp Asp lie val Asn Trp Leu Lys Lys

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TAT

110

103

ACA GCT GGC AGA GAG GCT GAT GAC ATC GTG AAC TGG CTG

8

80 (1)

Willows TYPE: other nucleic acid (semi-synthetic DNA)

SEQUENCE DESCRIPTION:

ATC AAG TGG GTT ACC TTC ATC TCT TTG TTG Met Lys Trp Val Thr Phe 11e Ser Leu Leu

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	776	Leu		636	Gla		GOG CIE GCG GCC CAC AAG		30				AAG	Gly Lys Lev Lys		ACG
	TIC	Phe		ည	Q A		GAG	Glu		TGG	Trp	5	999	Gly		228

TOC 1.3 1.4 (1) GTG Ser Lev Val TAC Val 000 S.S.D GXT HA.A.A. TTC 717 ACA 195 Gly Arg Asn 15. .T.T.C Asp Ile Pro GAC 929 CGG AAC AAC CAG Ile Lys His Asn Gln Leu Pro Leu Val ごの口 Ala Pro Lys Tie Phe Giy Gly Glu Ile Lys Thr His Ile Leu ATC AAA CAC SCA TTG Thr Ala Ala GOC GTG ATA S) H TCG CTC GAC AAA Leu Asp CCG 168 CCC MAG AGT Pro Lys CCA AS V 0%0 Glu Ser Ser Glu Val GBG 150 AST A.A.G 00 00 00 00 Phe Gly Ile Thr Ser TTT 501 TCC AGC GAG 83C TOT GOC 80.00 (2) (3) (3) TTT Lys asp Gly Val Val ATT Gla Ser GAG215 Ala Lys Gln Phe Leu Gln Ala Ala Glu 120 CAG ପ୍ରକ୍ରେ GRA Qi h 3.1.3 Glu Gly GAT GGG GTT 375 200 ACC MAG CIG ATC 636 58 188 GGA e A Ser Asp Try 3.03 3.03 3.14 CAG GTG CCC ACZ G1 u GAG 170 AAC GGT GAA ATC AAG GAC TAT Lys Gly Lys 250 Tay GTC GIC 300 TIT ALA GCT CTT 155 GGC AAG 10 (3) SIC ASO AAC 140 Val Les Phe Lys Lys Phe TTG CAG GCA CTC GAC ACC asp Giy Lys Leu Ser GTC Thr 220 ATC GGC Ile Gly Fhe Ser Asp Val Phe *GT 125 TIL 690 Lie ATC GAG Lys Glu Asa Leu ANG GAG AAC 17 37.0 205 GAC AAG AAG ACT STU 190 0.30 3.4.2 Les Fhe GTG GCA TTC 275 CAC Phe Thr Glu CIG TIC 777 in in रुपद D.X.Z. cre TTT TTC 330 260 ATC 200 32°C AGC 110 240 AAG 229 Lys Asp Leu TCC 23.0 GAT Ser Lys CIG GAG CTG ASP 145 232 AAC 1770 225 340 ATC GAA 110 CAG GAC A.A.A 130 CTG ASD 220 3.7°C TTC 510 00 01 01 554 808 750 702 846 798 894

30.5

Glu Phe Cys His Arg Phe Leu

Gin Gly

Lys Ile

Lys Pro His Lev

AZC.

AGC

CAG GAG

CCG

GAG GAC

TGG

G&C

AAG CAG CCT

GTC

AAG

STG

1134

335

330

325

Wet.

10.00

Leu 020

240

Glu Asp Trp

Asp Lys Gla Pro Val Lys Val

() +3

(3) 13)

999

AAG

MAC

TT

(A.A.

GAC

676

GAT GAG 350

4.4.4

A.A.B

A.A.C

1182

345

Led Val Gly Lys

det nio eus usa

12 A

(2) (3) GCT

Phe TTT

ASD

62.5

Lys Lys Asn

370

355

2.1.2

913

GAG

TTC TAT GCC

CCA

TGG

TGT GGT

CAC

Tac

450

1230

615 0.80

387 CTT

385

300

(a)

V&I

Glu she Tyr Ala Pro Trp Cys Gly His Cys Lys

981

CCC

ATT

706

MAM OTS

CCA

GAG

ACG. Tit

TAC AAG GAC CAT

GAG

1278

380

375

070

lle Trp

Asp GAT

Lys Leu Gly Glu

IÑ

5.57

Asp His Glu

330 AAC

800

(1) (2)

64 74 73

13 10 ATC

Val

130 PM MIG

285

Ser Thr TCG ACT

ALO.

Ash Glu Val

Giu

Val STC

910

STA

900 390

240

GAC

900

and eng

GTG

GAG

900 900

1325

35 275 Lys Lys Glu Glu Cys Pro Ala ATG Met ACA AAG 500 AGC 260 GAG The Lys Tyr Lys Pro Giu Ser Glu Glu ACC 3%0 Car 350 333 PFG TGC E TH 30 GAG TAC AAG 310 ACC TGC Tall 100 UN CAC CGC GAC AAC CCG 6CC dsa CCC GAA TCG 280 rs: 265 O.L. Val Arg Leu CAG CGC ATC GTG G)n CTG GAG CGC 210 33 13 5 GAG GAG CTC GGC Ile Zeu Gla 300 CTC Leu CTG 10 ATC 285 27.0 626 13. 13. ACC 17.7 acc acc TAT 270 Ala Glu Phe Phe Gly TIC TIT AAG GCA Leu CTG GAG a [5 GAC AGG 000 320 GAG 99 614 CAC Arg 305 Leu G124 GAG CTG CTG I2e MIC 1038 1086 942 990

70. 4. 8 8 8 8

2470 1518 5365 1422 2374 GAT (1) (1) (1) GAC A.A.G Lys GRI ASS. 2.50 ASD GAC der. Phe GAC Met Glu Glu Asp 228 Ala 4T.3 Phe Pro Ala Ser GGT Cly CAT Ala Gly Asp Asp GAC ATG GAG GAA AG3 000 CTG GAT Leu Asp 0 989 439 TTT CCT X VII SES. ACG. 010 Arg GAT GGG Gly Gle Asp Gly GAG CCA Phe CGC 277 460 610 Lys CTG CTC AAG GGG GAA Tyr Asn Gly Glu Lys Asp Glu Leu 610 GAA GGC CAG GCA GAG ner. 513 GAT ACA Thr AMC 425 GGT GAR gra AAA 333 Pro TAC 613 440 Ch ch ch \$4 00 00 323 Leu Glu GTG CIG GAG AGC Gln Lys Ala Val TIC Phe GAT G12 505 Q) (S) GTG CAC AGC ACG GTC ATT Leu Thr Val GAC Gir Asp (0) (1) CAG AAA 9104 GAG 277 127 3 4 5× 89 64 1.7.3 CSC 9

06.5

REQUENCE LENGTH: :ON GI Cas

SEQUENCE TYPE: amino

POPOLOGY: linear

TYPE: protein CLECULAR

Val Leu H. 13 DESCRIPTION SEQUENCE

Tyr 5 10 10 610 Tyr Leu Leu Val C

the Ala Glu Ala Leu Ala Ala Ris Lys 23

Arg Lys Val A.la Tyr Ale G1y Arg Leu Gle Z AT E Iys Ala Leu Ale Pro Gla Gla Gla A. B. Glu Gly Ser ren 3.00 Gin Ser 3.1 A Cys Gly His Cys Lys Gla ren Thr \$ V.15 C. संस 623 dog. 220 A.a. 50 413 Val 315

520 Ser A.28 ASE Val TUL 11.6 Gly Asp Asp Asn 350 Arg Ala 3.03 n to भ्राद Lys Fae Gly Arg 2.0 £13 13.0 Thr Thr 020 T\T 77x 37.5 G3.22 GIŽ 8 AT 10

Ala AJa G13 Asp Thr Leu Pro H K: Lys Arg Thr Gly Pro Ala Ala 320 5.7 m

GJ y Glu Val Ala Val Ile Ser Ser 135 gJn Val Giu Ser Leu Ala

160 614 Ala Ala Ser Ala Lys Gln Phe Leu Gln 11.2 11.3 Q SE 350 14 8) 14 G3n Lys Asp Val 145 Phe Ser Asn Ser Asp Val zuz 170 11e 775 Pro Phe 13.6 100 100 100 100 ASE (1), (6) Lie E C

Price 773 Lys 190 ಗಾನ್ Val Va. 615 33 Asp Lys Asp Len Gin 180 1×2 Lys

Asa n75 Lvs 242 Val Gla Asn the Glu Gly er. Gly Arg 375 Asp

Thr 200 GIS val Ile Leu 0 H Rt 3 His Asn Gln 1.73 Fhe Ile 193 ASD

215

A.5.

Sex

Lys

770

Leu

Val

Asp

sty ale pro Glu Giu Glu

240 Ile Lys G7n 235 GLy Lys Ile Phe Gly 230 Q1 Q1 Gln Thr Ala

Sex Asp Tyr Asp Gly Lys Lys Ser Val Ser 024 Ce. teu Leu Phe

Asn Phe Lys Thr Ala Ala Giu Ser Phe Lys Gly Lys Ile Leu Phe Ile Gly Leu Lys Lys Glu Glu Cys Pro Ala Val Arg Leu Ile Thr Leu Glu Phe Ile Asp Ser Asp His Thr Asp Asn Gln Arg Ile Leu Glu Phe Phe 305 Giu Glu Met Thr Lys Tyr Lys Pro Glu Ser Glu Glu Leu Thr Ala Glu 385 G1n Lys val Leu val Gly Lys Asn Phe Giu Asp Val Ala Phe Asp Glu Lys STH 27.0 Glu Asn Ile Val Ile Ala Lys Met Asp Ser Thr Ala Asn Glu Val Glu Lys Asn val the val Glu the Tyr Ala Pro Trp Cys Gly His Cys Lys The Lys Lys Phe Leu Glu Ser Gly Gly Gla Asp Gly Ala Gly Asp Asp Ala Val Lys Val His Ser Phe Pro Thr Leu Lys Phe Phe Pro Ala Ser He Thr Glu Phe Cys His Arg Phe Leu Glu Gly Lys Lie Lys Pro Leu Ala Pro Ile Trp Asp Lys Leu Gly Glu Thr Tyr Lys Asp His Lea Met Ser Gin Glu Lea Pro Gia Asp Trp Asp Lys Gin Pro Vai AS O 275 (3) (3) (3) Arg Thr Val Tle Asp Tyr Asn Gly Glu Arg Thr Leu Asp Gly 260 340 420 N 101 325 405 310 390 295 ري در در 280 350 4.20 265 W 150 423 250 330 619 (A) (A) (B) 395 300 285 365 270 350 255 11. 11. 320 400

> Asp Asp Asp Gin Lys Ala Val Asp Asp Leu Glu Asp Leu Glu Glu Ala Glu Glu Fro Asp Met Glu Glu \$70 Lys Asp Giu Leu 475 080

383

490

450

453

prief Description of the Drawings?

Fig. 1A illustrates a construction of the expression plasmid AHNFDILY1.

Fig. 18 illustrates a construction of the expression plasmid

Fig. 1C illustrates a construction of the expression plasmid

Fig. 2 shows a boundary of the HSA prepro-sequence and the PDI gene on a numan expression plasmid.

Fig. 3 is a photograph showing the result of SDS-polyacrylamide gel electrophoresis of an expressed and segreted crude recombinant numan PDI, wherein lane 1 is a molecular weight marker, lane 2 is pan/AH22 (control) and lane 3 is panhabilyl/AH22.

gig. 4 illustrates the separation of a recombinant human PDI by

lydrophobic column chromatography.

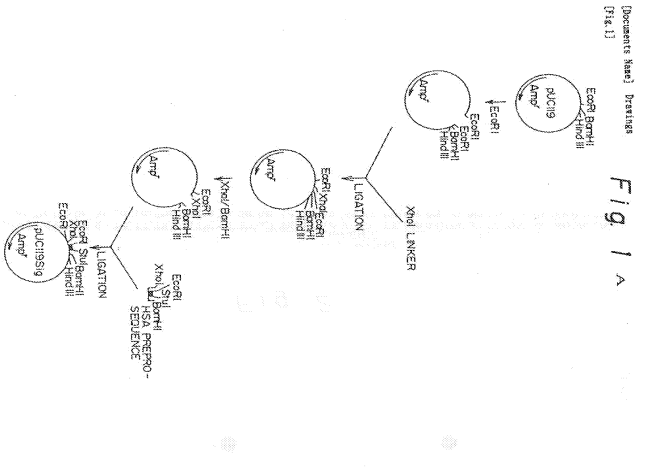
Fig. 5 shows the result of SDS-polyacrylamide gelalectrophoresis of a purified recombinant human PDI, wherein the numbers at the bottom correspond to the fraction numbers of the lydrophobic column chromatography shown in Fig. 4, and M is a molecular weight marker.

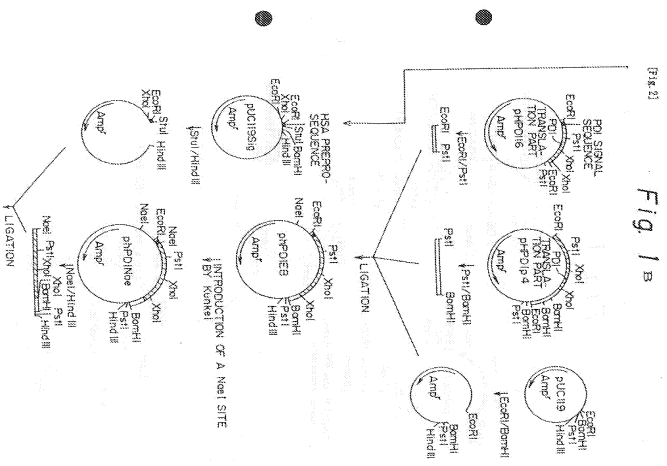
Fig. 6 is a photograph of SDS electrophoresis showing the axpression of human PDI in a yeast strain HIS13.

Fig. 7 is a photograph of SDS-polyacrylamide gel

numan PDI and HSR in the yeast strain HIS23

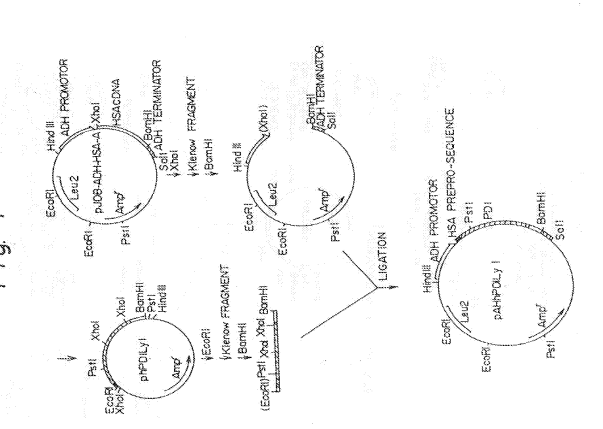
Fig. 8 shows the result of densitometric determination of the amount of secreted HSA using the SDS-polyacrylamide gel slectrophoresis gel in Fig. 7.





614

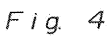
(Fig. 4)



(Fig. 33



Photo as substitute for drawing volecular 123
Weight 123
Secreted PDI
67,000
43,000
30,000
20,000



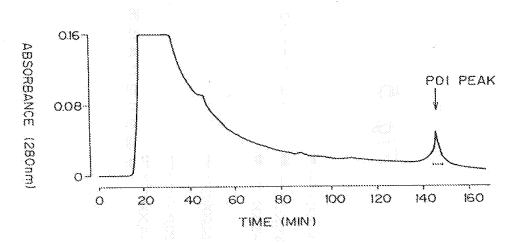


Fig. 6

(Fig. 83

Fig. 6

Photo as substitute for drawing

(Dalton)

94,000 - -

2: PAHAPOIL 91/HIS23 1: pAH/HIS23

[Fig. 7]

S

200029 43,000

94,000

10d

30,000

20,000

Photo as substitute for drawing

1, 3, 5, 7, 9: pAH/HIS23 2, 4, 6, 8, 10: pAHhPDILy1/HIS23

11: HSA STANDARD 0.25µg

AMOUNT OF SECRETED HSA {mg/1) 0 TRANSFORMANTS EXAMINED N Ċ) 4. (N Ø) ~1 α ω

1, 3, 5, 7, 9: pAH/HIS23 2, 4, 6, 8, 10: pAHhPDILy1/HIS23

ABSTRACT OF THE DISCLOSURE

abstract]

(Objects) It is an object of the present invention to provide an Ġ bject of the invention provides a co-expression of said gene and xpression of a protein disultide isomerase (PDI) gene. Another soreign gene cading for a useful polypeptide.

ene coding for the useful polypeptide, and then comexpressing their enes in the transformant. orocess for the preparation of PDI which comprises inserting a novel ontaining, in a co-expressible state, the fusion gene and a foreign libumin prepro sequence and a human PDI gene, transforming a host reparation of a useful polypeptide by preparing a transformant [Constitution] The present invention is characterized by a ension gene composed of a DNA fragment coding for human serum ell and then expressing the PDI gene; and a process for the

coordingly it is possible to improve a production effectency of the [Advantages] According to the present invention, a process for he large-scale production of the human PDI was established, and seful polypeptide by the above mentioned co-expression.

information of Applicant's History

Identification Number

(390022998)

1. Date of Change

16th November, 1990.

New Registration

(Reason for the Change)

1-1, Hitotsubashi 1-chome, chiyoda-ku, Tokyo.

Address

Name

TOMEN CORPORATION

Figures to be chosen] None